# H. R. 5629

To amend the Public Health Service Act to establish a pathway for the licensure of biosimilar biological products, and for other purposes.

## IN THE HOUSE OF REPRESENTATIVES

March 13, 2008

Ms. Eshoo (for herself, Mr. Barton of Texas, Mr. Hill, Mr. Upton, Mr. Wynn, Mr. Pitts, Ms. Zoe Lofgren of California, Mr. Rogers of Michigan, Mr. Capuano, Mr. Buyer, Mr. McGovern, Mr. Tim Murphy of Pennsylvania, Mr. Lynch, and Mr. Ferguson) introduced the following bill; which was referred to the Committee on Energy and Commerce, and in addition to the Committee on the Judiciary, for a period to be subsequently determined by the Speaker, in each case for consideration of such provisions as fall within the jurisdiction of the committee concerned

# A BILL

To amend the Public Health Service Act to establish a pathway for the licensure of biosimilar biological products, and for other purposes.

- 1 Be it enacted by the Senate and House of Representa-
- 2 tives of the United States of America in Congress assembled,
- 3 SECTION 1. SHORT TITLE.
- 4 This Act may be cited as the "Pathway for
- 5 Biosimilars Act".

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#### TITLE I—AMENDMENTS TO PUBLIC HEALTH SERVICE ACT

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# 3 TITLE I—AMENDMENTS TO

# 4 PUBLIC HEALTH SERVICE ACT

### 5 SEC. 101. LICENSURE PATHWAY FOR BIOSIMILAR BIOLOGI-

- 6 CAL PRODUCTS.
- 7 (a) Licensure of Biological Products as Bio-
- 8 SIMILAR OR INTERCHANGEABLE.—Section 351 of the
- 9 Public Health Service Act (42 U.S.C. 262) is amended—
- 10 (1) in subsection (a)(1)(A), by inserting "under
- this subsection or subsection (k)" after "biologics li-
- cense"; and
- (2) by adding at the end the following:
- 14 "(k) Licensure of Biological Products as Bio-
- 15 SIMILAR.—
- 16 "(1) IN GENERAL.—Any person may submit an
- 17 application for licensure of a biological product
- under this subsection.
- 19 "(2) CONTENT.—

1	"(A) REQUIRED INFORMATION.—An appli-
2	cation submitted under this subsection shall in-
3	clude information demonstrating that—
4	"(i) the biological product is bio-
5	similar to a reference product based upon
6	data derived from—
7	"(I) analytical studies that dem-
8	onstrate that the biological product is
9	highly similar to the reference product
10	notwithstanding minor differences in
11	clinically inactive components;
12	"(II) animal studies (including
13	the assessment of toxicity); and
14	"(III) a clinical study or studies
15	(including, but not limited to, the as-
16	sessment of immunogenicity and phar-
17	macokinetics or pharmacodynamics)
18	that are sufficient to demonstrate
19	safety, purity, and potency for each
20	condition of use for which the ref-
21	erence product is approved;
22	"(ii) the biological product and ref-
23	erence product utilize the same mechanism
24	or mechanisms of action for the condition
25	or conditions of use prescribed, rec-

1	ommended, or suggested in the proposed
2	labeling, but only to the extent the mecha-
3	nism or mechanisms of action are known
4	for the reference product;
5	"(iii) the condition or conditions of
6	use prescribed, recommended, or suggested
7	in the labeling proposed for the biological
8	product have been previously approved for
9	the reference product;
10	"(iv) the route of administration, the
11	dosage form, and the strength of the bio-
12	logical product are the same as those of
13	the reference product; and
14	"(v) the facility in which the biological
15	product is manufactured, processed,
16	packed, or held meets standards designed
17	to assure that the biological product con-
18	tinues to be safe, pure, and potent.
19	"(B) Waiver regarding analytical
20	STUDIES, ANIMAL STUDIES, AND CLINICAL
21	STUDIES.—
22	"(i) IN GENERAL.—The Secretary
23	may, in the Secretary's discretion, deter-
24	mine that an element described in sub-
25	clause (I), (II), or (III) of subparagraph

1	(A)(i) is unnecessary and waive the re-
2	quirement that such element be submitted
3	in an application under this subsection.
4	"(ii) Assessments of
5	IMMUNOGENICITY.—Notwithstanding
6	clause (i), the Secretary may determine
7	that an assessment of immunogenity de-
8	scribed in subparagraph (A)(i)(III) is un-
9	necessary and waive the requirement that
10	such an assessment be submitted in an ap-
11	plication under this subsection only if the
12	Secretary has published a final guidance,
13	following receipt and consideration of pub-
14	lic comments on a draft guidance—
15	"(I) advising that it is feasible in
16	the current state of scientific knowl-
17	edge to make determinations on
18	immunogenicity with respect to prod-
19	ucts in the product class to which the
20	biological product belongs; and
21	"(II) explaining the data that
22	will be required to support such a de-
23	termination.
24	"(C) Additional information.—An ap-
25	plication submitted under this subsection—

1	"(i) shall include publicly-available in-
2	formation regarding the Secretary's pre-
3	vious determination that the reference
4	product is safe, pure, and potent; and
5	"(ii) may include any additional infor-
6	mation in support of the application, in-
7	cluding publicly-available information with
8	respect to the reference product or another
9	biological product.
10	"(3) Evaluation by secretary.—Upon re-
11	view of an application (or a supplement to an appli-
12	cation) submitted under this subsection, the Sec-
13	retary shall approve the application (or the supple-
14	ment) if—
15	"(A) the Secretary determines that the in-
16	formation submitted in the application (or the
17	supplement) is sufficient to show that the bio-
18	logical product is biosimilar to the reference
19	product with respect to each condition of use
20	for which the reference product is approved;
21	and
22	"(B) the applicant (or other appropriate
23	person) consents to the inspection of the facility
24	that is the subject of the application, in accord-
25	ance with subsection (c).

1	"(4) Safety standards for determining
2	INTERCHANGEABILITY.—
3	"(A) Determination.—Upon review of
4	an application submitted under this subsection
5	or any supplement to such application, the Sec-
6	retary shall determine the biological product to
7	be interchangeable with the reference product if
8	the Secretary determines that the information
9	submitted in the application (or a supplement
10	to such application) is sufficient to show that—
11	"(i) the biological product—
12	"(I) is biosimilar to the reference
13	product and any biological product li-
14	censed under this subsection that has
15	been determined to be interchangeable
16	with the reference product; and
17	"(II) can be expected to produce
18	the same clinical result as the ref-
19	erence product in any given patient
20	for each condition of use prescribed,
21	recommended, or suggested in the la-
22	beling of the reference product; and
23	"(ii) for a biological product that is
24	administered more than once to an indi-
25	vidual, the risk in terms of safety or dimin-

1	ished efficacy of alternating or switching
2	between use of the biological product and
3	the reference product is not greater than
4	the risk of using the reference product
5	without such alternation or switch.
6	"(B) Guidelines.—Notwithstanding sub-
7	paragraph (A), the Secretary may not make a
8	determination that a biological product licensed
9	under this subsection is interchangeable with
10	the reference product unless the Secretary has
11	published a final guidance, following receipt and
12	consideration of public comments on a draft
13	guidance—
14	"(i) advising that it is feasible in the
15	current state of scientific knowledge to
16	make such determinations with respect to
17	products in the product class to which that
18	biological product belongs; and
19	"(ii) explaining the data that will be
20	required to support such a determination
21	"(5) General rules.—
22	"(A) ONE REFERENCE PRODUCT PER AP-
23	PLICATION.—A biological product, in an appli-
24	cation submitted under this subsection, may not

be evaluated against more than 1 reference
 product.

- "(B) Review.—An application submitted under this subsection shall be reviewed by the division within the Food and Drug Administration that is responsible for the review and approval of the application under which the reference product is licensed.
- "(C) RISK EVALUATION AND MITIGATION STRATEGIES.—The authority of the Secretary with respect to risk evaluation and mitigation strategies under the Federal Food, Drug, and Cosmetic Act shall apply to biological products licensed under this subsection in the same manner as such authority applies to biological products licensed under subsection (a).
- "(D) LISTED SELECT AGENTS AND TOX-INS.—If information in an application submitted under this subsection, in a supplement to such an application, or otherwise available to the Secretary shows that a biological product is, bears, or contains a select agent or toxin listed in section 73.3 or 73.4 of title 42, section 121.3 or 121.4 of title 9, or section 331.3 of title 7 of the Code of Federal Regulations (or any suc-

1	cessor regulations), the Secretary shall not li-
2	cense the biological product under this sub-
3	section.
4	"(6) Exclusivity for first interchange-
5	ABLE BIOLOGICAL PRODUCT.—The Secretary shall
6	not make a determination under paragraph (4) that
7	a second or subsequent biological product is inter-
8	changeable with the same reference product for
9	which a prior biological product has received a deter-
10	mination of interchangeability until 24 months after
11	the later of—
12	"(A) the date of the first commercial mar-
13	keting of the first biosimilar biological product
14	determined to be interchangeable for that ref-
15	erence product; or
16	"(B) with respect to a product marketed
17	before the date the product is determined to be
18	interchangeable, the date that the product is
19	determined to be interchangeable.
20	"(7) Exclusivity for reference prod-
21	UCT.—
22	"(A) EFFECTIVE DATE OF BIOSIMILAR AP-
23	PLICATION LICENSURE.—Subject to subpara-
24	graph (D) and paragraph (8), approval of an
25	application under this subsection may not be

1	made effective by the Secretary until the date
2	that is 12 years after the date on which the ref-
3	erence product was first licensed under sub-
4	section (a).
5	"(B) FILING PERIOD.—An application
6	under this subsection may not be submitted to
7	the Secretary until the later of—
8	"(i) the date of commencement of a
9	proceeding for issuance of guidance pursu-
10	ant to paragraph (9) with respect to the
11	product class within which the product
12	that is the subject of such application falls;
13	or
14	"(ii) the date that is 4 years after the
15	date on which the reference product was
16	first licensed under subsection (a).
17	"(C) First licensure.—For purposes of
18	this paragraph, the date on which the reference
19	product was first licensed under subsection (a)
20	does not include the date of approval of a sup-
21	plement or of a subsequent application for a
22	new indication, route of administration, dosage
23	form, or strength for the previously licensed ref-
24	erence product.

"(D) Medically Significant New Indication.—If, during the 8-year period following licensure of the reference product, the Secretary approves a supplement to the application for the reference product that seeks approval to market the reference product for a new indication that, if approved, would be a significant improvement, compared to marketed products, in the treatment, diagnosis, or prevention of disease, approval of an application submitted under this subsection may not be made effective by the Secretary until the date that is 14 years after the date on which the reference product was first licensed under subsection (a).

# "(8) Pediatric studies.—

"(A) EXCLUSIVITY.—If, before or after licensure of the reference product under subsection (a) of this section, the Secretary determines that information relating to the use of such product in the pediatric population may produce health benefits in that population, the Secretary makes a written request for pediatric studies (which shall include a timeframe for completing such studies), the applicant or holder of the approved application agrees to the re-

1	quest, such studies are completed using appro-
2	priate formulations for each age group for
3	which the study is requested within any such
4	timeframe, and the reports thereof are sub-
5	mitted and accepted in accordance with section
6	505A(d)(3) of the Federal Food, Drug, and
7	Cosmetic Act—
8	"(i) the period referred to in para-
9	graph (7)(A) of this subsection is deemed
10	to be 12 years and 6 months rather than
11	12 years; and
12	"(ii) if paragraph (7)(D) of this sub-
13	section applies, the period referred to in
14	such paragraph is deemed to be 14 years
15	and 6 months rather than 14 years.
16	"(B) Exception.—The Secretary shall
17	not extend the period referred to in subpara-
18	graph (A)(i) or (A)(ii) of this paragraph if the
19	determination under section 505A(d)(3) of the
20	Federal Food, Drug, and Cosmetic Act is made
21	later than 9 months prior to the expiration of
22	such period.
23	"(C) Application of Certain Provi-
24	SIONS.—The provisions of subsections (a), (d),
25	(e), (f), (h), (j), (k), and (l) of section 505A of

the Federal Food, Drug, and Cosmetic Act shall apply with respect to the extension of a period under subparagraph (A) of this paragraph to the same extent and in the same manner as such provisions apply with respect to the extension of a period under subsection (b) or (c) of section 505A of the Federal Food, Drug, and Cosmetic Act.

# "(9) Guidance documents.—

"(A) IN GENERAL.—The Secretary shall, after opportunity for public comment, issue final guidance with respect to the licensure under this subsection of a biological product or product class. Such guidance shall be issued in accordance, except as provided in subparagraph (B)(i), with section 701(h) of the Federal Food, Drug, and Cosmetic Act.

# "(B) Public comment.—

"(i) IN GENERAL.—Before issuing final guidance under subparagraph (A), the Secretary shall publish a proposed guidance, provide an opportunity for the public to comment on the proposed guidance, and publish a response to comments received under this clause.

1	"(ii) Input regarding most valu-
2	ABLE GUIDANCE.—The Secretary shall es-
3	tablish a process through which the public
4	may provide the Secretary with input re-
5	garding priorities for issuing guidance.
6	"(C) CERTAIN PRODUCT CLASSES.—
7	"(i) Guidance.—The Secretary may
8	indicate in a guidance document under
9	subparagraph (A) that the Secretary will
10	not license a product or product class (not
11	including any recombinant protein) under
12	this subsection because the science and ex-
13	perience, as of the date of such guidance,
14	does not allow such licensure.
15	"(ii) Modification or reversal.—
16	The Secretary may issue a subsequent
17	guidance document under subparagraph
18	(A) to modify or reverse a guidance docu-
19	ment under clause (i).
20	"(D) PETITION FOR INITIATION OF GUID-
21	ANCE FOR CERTAIN PRODUCTS.—In the case of
22	a reference product that was licensed by the
23	Secretary more than 7 years prior to the date
24	of the enactment of the Pathway for
25	Biosimilars Act, a person may petition the Sec-

retary at any time to commence the process for issuing final guidance under subparagraph (A) for the product class to which the reference product belongs. Any such petition shall include a description of the scientific feasibility and rationale for the request. For guidance petitioned under this subparagraph, the Secretary shall, within 2 years of such petition, issue final guidance with respect to that product class.

"(E) Requirement for application consideration.—The Secretary may not accept an application under this subsection until the Secretary has initiated a proceeding for issuance of guidance with respect to the product class within which the product that is the subject of the application falls. The Secretary may not approve an application under this subsection until the Secretary has completed the proceeding for issuance of guidance with respect to the product class within which the product that is the subject of the application falls.

"(F) REQUIREMENT FOR PRODUCT CLASS-SPECIFIC GUIDANCE.—Product class-specific

1	guidance issued under subparagraph (A) shall
2	include a description of—
3	"(i) the criteria that the Secretary will
4	use to determine whether a biological prod-
5	uct is biosimilar to a reference product in
6	such product class;
7	"(ii) the criteria, if available, that the
8	Secretary will use to determine whether a
9	biological product meets the standards for
10	interchangeability described in paragraph
11	(4); and
12	"(iii) the criteria, if available, that the
13	Secretary will use to assess
14	immunogenicity.
15	"(10) Naming.—The Secretary shall ensure
16	that the labeling and packaging of each biological
17	product licensed under this subsection bears a name
18	that uniquely identifies the biological product and
19	distinguishes it from the reference product and any
20	other biological products licensed under this sub-
21	section following evaluation against such reference
22	product.
23	"(1) PATENT NOTICES; RELATIONSHIP TO FINAL AP-
24	PROVAL.—

1	"(1) Definitions.—For the purposes of this
2	subsection, the term—
3	"(A) 'biosimilar product' means the bio-
4	logical product that is the subject of the appli-
5	cation under subsection (k);
6	"(B) 'relevant patent' means a patent
7	that—
8	"(i) expires after the date specified in
9	subsection $(k)(7)(A)$ that applies to the
10	reference product; and
11	"(ii) could reasonably be asserted
12	against the applicant due to the unauthor-
13	ized making, use, sale, or offer for sale
14	within the United States, or the importa-
15	tion into the United States of the bio-
16	similar product, or materials used in the
17	manufacture of the biosimilar product, or
18	due to a use of the biosimilar product in
19	a method of treatment that is indicated in
20	the application;
21	"(C) 'reference product sponsor' means the
22	holder of an approved application or license for
23	the reference product; and
24	"(D) 'interested third party' means a per-
25	son other than the reference product sponsor

that owns a relevant patent, or has the right to commence or participate in an action for infringement of a relevant patent.

> "(2) Handling of confidential informa-TION.—Any entity receiving confidential information pursuant to this subsection shall designate one or more individuals to receive such information. Each individual so designated shall execute an agreement in accordance with regulations promulgated by the Secretary. The regulations shall require each such individual to take reasonable steps to maintain the confidentiality of information received pursuant to this subsection and use the information solely for purposes authorized by this subsection. The obligations imposed on an individual who has received confidential information pursuant to this subsection shall continue until the individual returns or destroys the confidential information, a court imposes a protective order that governs the use or handling of the confidential information, or the party providing the confidential information agrees to other terms or conditions regarding the handling or use of the confidential information.

> "(3) Public Notice by Secretary.—Within 30 days of acceptance by the Secretary of an appli-

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1	cation filed under subsection (k), the Secretary shall
2	publish a notice identifying—
3	"(A) the reference product identified in the
4	application; and
5	"(B) the name and address of an agent
6	designated by the applicant to receive notices
7	pursuant to paragraph (4)(B).
8	"(4) Exchanges concerning patents.—
9	"(A) EXCHANGES WITH REFERENCE
10	PRODUCT SPONSOR.—
11	"(i) Within 30 days of the date of ac-
12	ceptance of the application by the Sec-
13	retary, the applicant shall provide the ref-
14	erence product sponsor with a copy of the
15	application and information concerning the
16	biosimilar product and its production. This
17	information shall include a detailed de-
18	scription of the biosimilar product, its
19	method of manufacture, and the materials
20	used in the manufacture of the product.
21	"(ii) Within 60 days of the date of re-
22	ceipt of the information required to be pro-
23	vided under clause (i), the reference prod-
24	uct sponsor shall provide to the applicant
25	a list of relevant patents owned by the ref-

erence product sponsor, or in respect of
which the reference product sponsor has
the right to commence an action of infringement or otherwise has an interest in
the patent as such patent concerns the biosimilar product.

"(iii) If the reference product sponsor

"(iii) If the reference product sponsor is issued or acquires an interest in a relevant patent after the date on which the reference product sponsor provides the list required by clause (ii) to the applicant, the reference product sponsor shall identify that patent to the applicant within 30 days of the date of issue of the patent, or the date of acquisition of the interest in the patent, as applicable.

# "(B) Exchanges with interested third parties.—

"(i) At any time after the date on which the Secretary publishes a notice for an application under paragraph (3), any interested third party may provide notice to the designated agent of the applicant that the interested third party owns or has rights under 1 or more patents that may

1 be relevant patents. The notice shall iden-2 tify at least 1 patent and shall designate 3 an individual who has executed an agreement in accordance with paragraph (2) to receive confidential information from the 6 applicant. 7 "(ii) Within 30 days of the date of re-8 ceiving notice pursuant to clause (i), the 9 applicant shall send to the individual des-10 ignated by the interested third party the 11 information specified in subparagraph 12 (A)(i), unless the applicant and interested 13 third party otherwise agree. "(iii) Within 90 days of the date of 14 15 receiving information pursuant to clause 16 (ii), the interested third party shall provide 17 to the applicant a list of relevant patents 18 which the interested third party owns, or 19 in respect of which the interested third 20 party has the right to commence or partici-21 pate in an action for infringement. 22 "(iv) If the interested third party is 23 issued or acquires an interest in a relevant 24 patent after the date on which the inter-

ested third party provides the list required

1	by clause (iii), the interested third party
2	shall identify that patent within 30 days of
3	the date of issue of the patent, or the date
4	of acquisition of the interest in the patent,
5	as applicable.
6	"(C) Identification of basis for in-
7	FRINGEMENT.—For any patent identified under
8	clause (ii) or (iii) of subparagraph (A) or under
9	clause (iii) or (iv) of subparagraph (B), the ref-
10	erence product sponsor or the interested third
11	party, as applicable—
12	"(i) shall explain in writing why the
13	sponsor or the interested third party be-
14	lieves the relevant patent would be in-
15	fringed by the making, use, sale, or offer
16	for sale within the United States, or im-
17	portation into the United States, of the
18	biosimilar product or by a use of the bio-
19	similar product in treatment that is indi-
20	cated in the application;
21	"(ii) may specify whether the relevant
22	patent is available for licensing; and
23	"(iii) shall specify the number and
24	date of expiration of the relevant patent.

1	"(D) CERTIFICATION BY APPLICANT CON-
2	CERNING IDENTIFIED RELEVANT PATENTS.—
3	Not later than 45 days after the date on which
4	a patent is identified under clause (ii) or (iii) of
5	subparagraph (A) or under clause (iii) or (iv) of
6	subparagraph (B), the applicant shall send a
7	written statement regarding each identified pat-
8	ent to the party that identified the patent. Such
9	statement shall either—
10	"(i) state that the applicant will not
11	commence marketing of the biosimilar
12	product and has requested the Secretary to
13	not grant final approval of the application
14	before the date of expiration of the noticed
15	patent; or
16	"(ii) provide a detailed written expla-
17	nation setting forth the reasons why the
18	applicant believes—
19	"(I) the making, use, sale, or
20	offer for sale within the United
21	States, or the importation into the
22	United States, of the biosimilar prod-
23	uct, or the use of the biosimilar prod-
24	uct in a treatment indicated in the ap-

1	plication,	would	not	infringe	the	pat-
2	ent; or					

3 "(II) the patent is invalid or un-4 enforceable.

> "(5) ACTION FOR INFRINGEMENT INVOLVING REFERENCE PRODUCT SPONSOR.—If an action for infringement concerning a relevant patent identified by the reference product sponsor under clause (ii) or (iii) of paragraph (4)(A), or by an interested third party under clause (iii) or (iv) of paragraph (4)(B), is brought within 60 days of the date of receipt of a statement under paragraph (4)(D)(ii), and the court in which such action has been commenced determines the patent is infringed prior to the date applicable under subsection (k)(7)(A), (k)(7)(D), or (k)(8) the Secretary shall make approval of the application effective on the day after the date of expiration of the patent that has been found to be infringed. If more than one such patent is found to be infringed by the court, the approval of the application shall be made effective on the day after the date that the last such patent expires.

> "(6) Limitations on actions for declaratory judgment.—With respect to a patent that is the subject of an explanation under paragraph

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1	(4)(D)(ii), no action for a declaratory judgment that
2	the patent is invalid, unenforceable, or not infringed
3	may be brought under section 2201 of title 28,
4	United States Code, by an applicant prior to the
5	date that is the later of—
6	"(A) 3 years prior to the date applicable
7	under subsection (k)(7)(A); or
8	"(B) 120 days after such explanation has
9	been provided.".
10	(b) Products Previously Approved Under Sec-
11	TION 505.—
12	(1) REQUIREMENT TO FOLLOW SECTION 351.—
13	Except as provided in paragraph (2), an application
14	for a biological product shall be submitted under
15	section 351 of the Public Health Service Act (42
16	U.S.C. 262) (as amended by this Act).
17	(2) Exception.—An application for a biologi-
18	cal product may be submitted under section 505 of
19	the Federal Food, Drug, and Cosmetic Act (21
20	U.S.C. 355) if—
21	(A) such biological product is in a product
22	class for which a biological product in such
23	product class is the subject of an application
24	approved under such section 505 not later than
25	the date of enactment of this Act; and

1	(B) such application—
2	(i) has been submitted to the Sec-
3	retary of Health and Human Services (re-
4	ferred to in this Act as the "Secretary")
5	before the date of enactment of this Act;
6	or
7	(ii) is submitted to the Secretary not
8	later than the date that is 10 years after
9	the date of enactment of this Act.
10	(3) Limitation.—Notwithstanding paragraph
11	(2), an application for a biological product may not
12	be submitted under section 505 of the Federal Food,
13	Drug, and Cosmetic Act (21 U.S.C. 355) if there is
14	another biological product approved under sub-
15	section (a) of section 351 of the Public Health Serv-
16	ice Act that could be a reference product with re-
17	spect to such application (within the meaning of
18	such section 351) if such application were submitted
19	under subsection (k) of such section 351.
20	(4) Deemed approved under section 351.—
21	An approved application for a biological product
22	under section 505 of the Federal Food, Drug, and
23	Cosmetic Act (21 U.S.C. 355) shall be deemed to be

a license for the biological product under such sec-

1	tion 351 on the date that is 10 years after the date
2	of enactment of this Act.
3	(5) Definitions.—For purposes of this sub-
4	section, the term "biological product" has the mean-
5	ing given such term under section 351 of the Public
6	Health Service Act (42 U.S.C. 262) (as amended by
7	this Act).
8	SEC. 102. FEES RELATING TO BIOSIMILAR BIOLOGICAL
9	PRODUCTS.
10	Subparagraph (B) of section 735(1) of the Federal
11	Food, Drug, and Cosmetic Act (21 U.S.C. 379g(1)) is
12	amended by inserting ", including licensure of a biological
13	product under section 351(k) of such Act" before the pe-
14	riod at the end.
15	TITLE II—AMENDMENTS TO
16	PATENT ACT
17	SEC. 201. AMENDMENTS TO CERTAIN PATENT PROVISIONS.
18	Section 271(e)(2) of title 35, United States Code is
19	amended—
20	(1) in subparagraph (A), by striking "or" after
21	"patent";
22	(2) in subparagraph (B), by adding "or" after
23	the comma at the end; and
24	(3) by inserting the following after subpara-
25	graph (B):

1	"(C)	a	$\operatorname{st}$	atem	nent	under	section
2	351(l)(4)(D	)(ii)	of	the	Public	Health	Service
3	Act,".						

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